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(54) **CHARGING CONTROL METHOD FOR SECONDARY CELL AND CHARGING CONTROL DEVICE FOR SECONDARY CELL**

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See application file for complete search history.

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H01M 4/505 (2010.01)

H01M 4/525 (2010.01)

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CPC **H02J 7/007** (2013.01); **H01M 4/505** (2013.01); **H01M 4/525** (2013.01); **H01M 10/44** (2013.01); **H02J 7/008** (2013.01); **H01M 10/052** (2013.01); **Y02E 60/122** (2013.01)

(58) **Field of Classification Search**

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(57) **ABSTRACT**

A method for controlling charging of a secondary cell including a positive electrode containing a positive-electrode active substance having that increases resistance in accordance with an increase in the SOC, a negative electrode; and a non-aqueous electrolyte includes performing constant-current charging at a set current value to a prescribed upper-limit voltage, performing constant-voltage charging at the upper-limit voltage after the upper-limit voltage V_1 has been reached, and terminating charging of the secondary cell when the charging current in the constant-voltage charging has decreased to a cutoff current value, the cutoff current value being set to a current value that complies with the relationships in formulas (I) and (II):

$$\text{Cutoff current value } A_2 \geq \text{set current value } A_1 \times X \quad (\text{I})$$

$$X = (\text{cell resistance } R_1 \text{ of secondary cell in target SOC } [\Omega] \times \text{set current value } A_1 [\text{A}]) / \text{upper-limit voltage } V_1 [\text{V}]. \quad (\text{II})$$

5 Claims, 5 Drawing Sheets

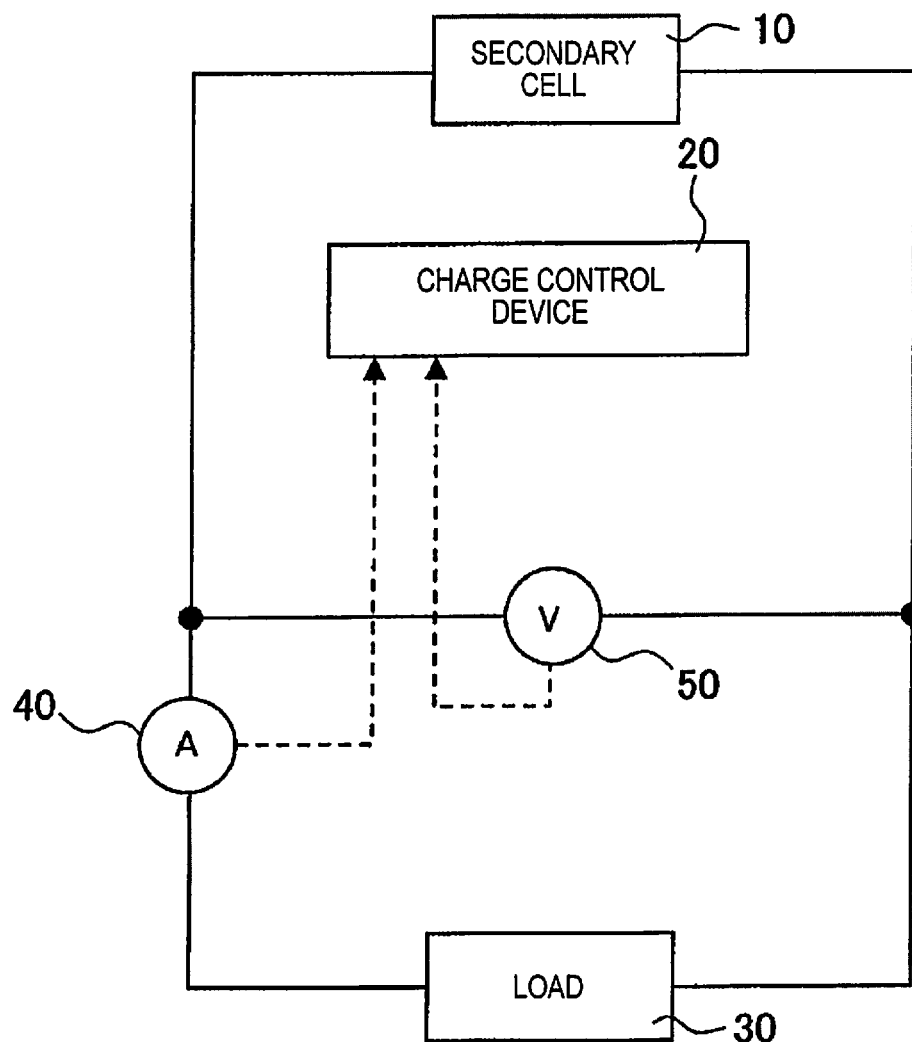


FIG. 1

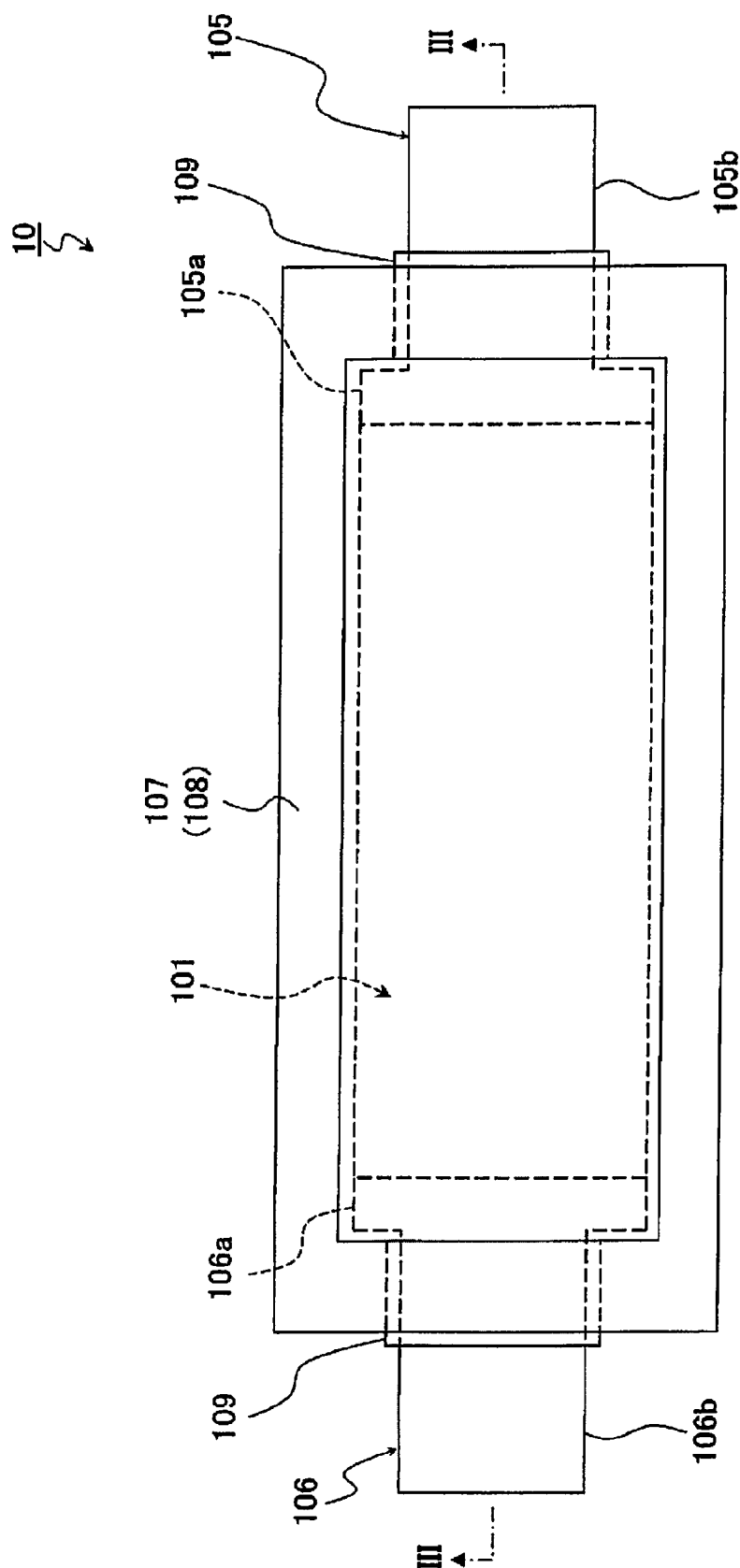


FIG. 2

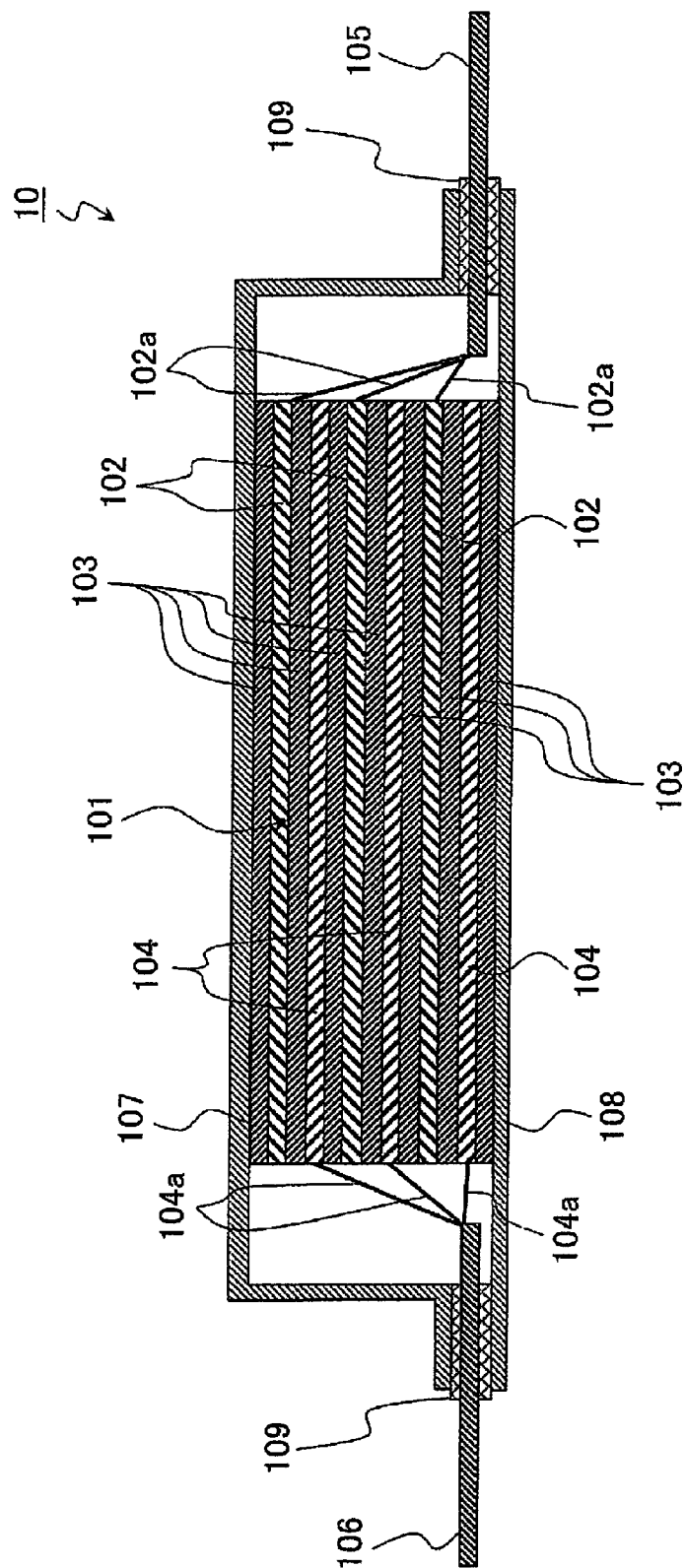


FIG. 3

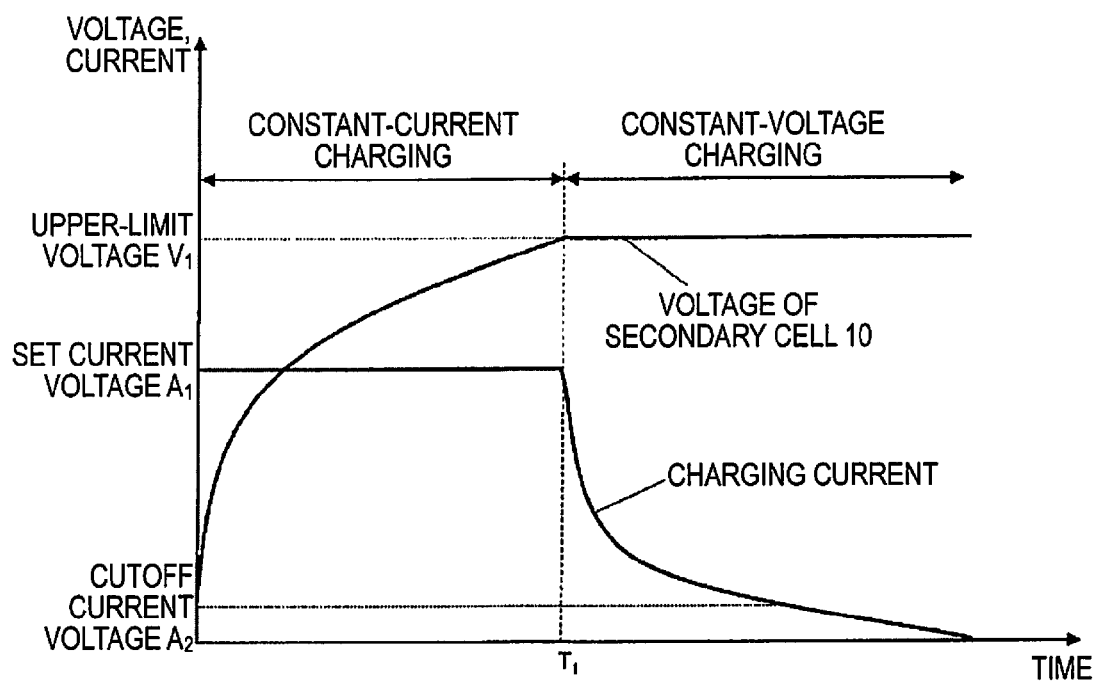


FIG. 4

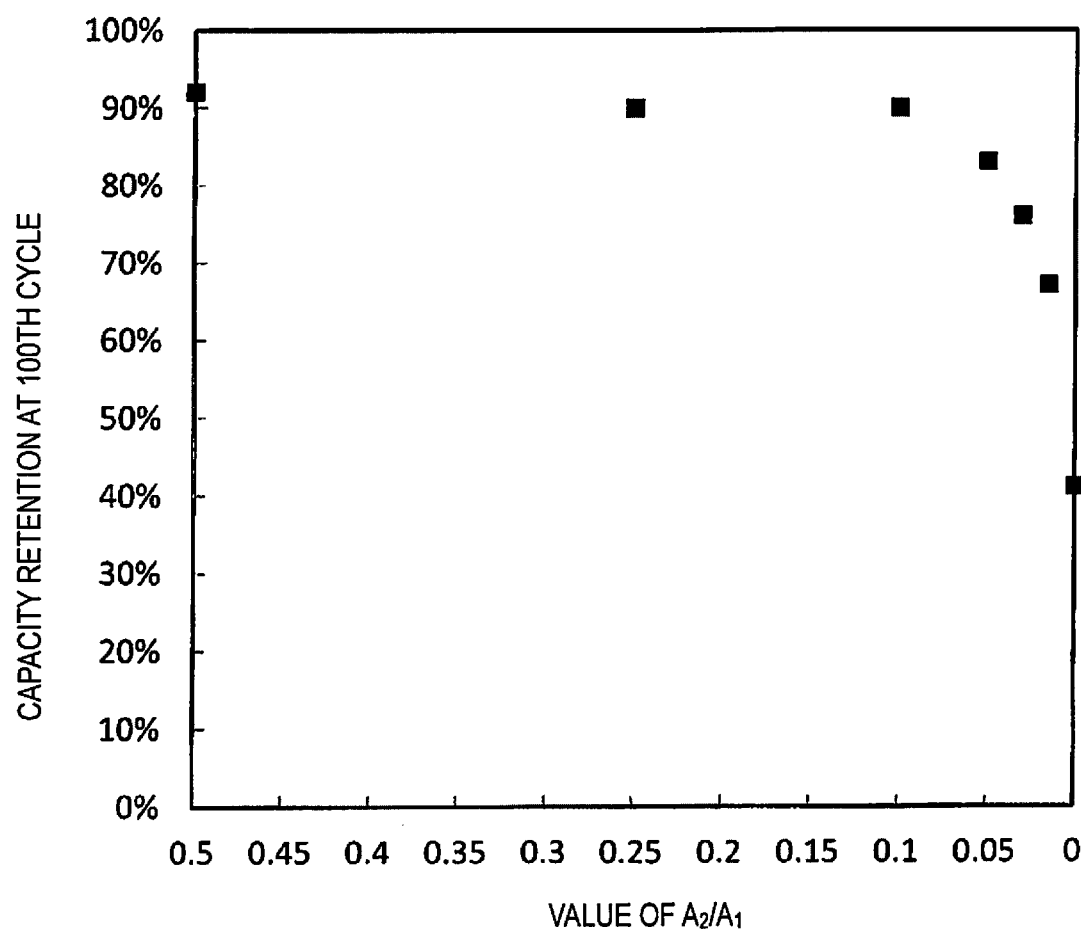


FIG. 5

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CHARGING CONTROL METHOD FOR SECONDARY CELL AND CHARGING CONTROL DEVICE FOR SECONDARY CELL

CROSS-REFERENCE TO RELATED APPLICATIONS

This application is a U.S. National stage application of International Application No. PCT/JP2013/064182 filed, May 22, 2013, which claims priority to Japanese Patent Application No. 2012-156668 filed in Japan on Jul. 12, 2012, the contents of each of which are hereby incorporated herein by reference.

BACKGROUND

1. Field of the Invention

The present invention relates to a method and device for controlling charging of a secondary cell.

2. Background Information

In recent years, a variety of positive-electrode active substances have been examined with the objective of raising the voltage and capacity of lithium-ion secondary cells and other secondary cells. Such positive-electrode active substances include, e.g., Li_2MnO_3 — LiMO_2 (where M is a transition metal having an average oxidation state of +3) and other solid solution materials disclosed in Japanese Laid-Open Patent Application No. 2008-270201.

SUMMARY

A feature of positive-electrode materials having high theoretical capacity, such as the solid solution materials disclosed in Japanese Laid-Open Patent Application No. 2008-270201, is that resistance increases in a high-SOC region. Therefore, when a secondary cell using such positive-electrode materials is charged using constant-current/constant-voltage charging, the positive/negative electrode potential changes during the constant-voltage charging process (particularly in the end stage of charging), and non-aqueous-electrolyte decomposition, lithium deposition, and other side reactions occur, resulting in diminished cycling characteristics.

The present invention improves the cycling characteristics of a secondary cell in which there is used, as a positive-electrode material, a positive-electrode active substance having the feature that resistance increases in accordance with an increase in the SOC.

The present invention solves the aforementioned problem by a method for controlling charging of a secondary cell in which there is used, as a positive-electrode material, a positive-electrode active substance having an open-circuit voltage curve that differs during charging and during discharging, wherein a cutoff current value A_2 for charging using constant-current/constant-voltage charging is set to a current value equal to or greater than the product ($A_1 \times X$) of a set current value A_1 during constant-current charging and a threshold coefficient X calculated from a cell resistance R_1 of a secondary cell at a target SOC, the set current value A_1 during constant-current charging, and the upper-limit voltage of charging.

According to the present invention, charging is terminated at a prescribed cutoff current when charging a secondary cell in which there is used, as a positive-electrode material, a positive-electrode active substance having the feature that resistance increases in accordance with an increase in the SOC, whereby side reactions associated with excessive current impressing can be suppressed. This makes it possible to

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improve the charge/discharge efficiency of the secondary cell, resulting in an improvement in the cycling characteristics.

BRIEF DESCRIPTION OF THE DRAWINGS

Referring now to the attached drawings which form a part of this original disclosure.

FIG. 1 is a schematic overview showing the charge control system for a secondary cell of the present embodiment.

FIG. 2 is a plan view of the secondary cell of the present embodiment.

FIG. 3 is a sectional view of the secondary cell across line in FIG. 2.

FIG. 4 is a profile showing the charging current and the change in voltage of the secondary cell of the present embodiment when the secondary cell is charged using constant-current/constant-voltage charging.

FIG. 5 is a graph showing the relationship between the “cutoff current value A_2 /set current value A_1 ” value obtained in the working and comparative examples, and the capacity retention at the 100th cycle.

DETAILED DESCRIPTION OF EMBODIMENTS

A preferred embodiment of the present invention is described below based on the accompanying drawings.

FIG. 1 is a view of the configuration of a system for controlling charging of the secondary cell of the present embodiment. This charge control system comprises a secondary cell 10, a control device 20, a load 30, an ammeter 40, and a voltmeter 50, as shown in FIG. 1.

The charge control device 20 is used for controlling the secondary cell 10, the charge control device 20 performing a control for charging and discharging the secondary cell 10 based on the charging/discharging current flowing through the secondary cell 10 as detected by the ammeter 40 and the terminal voltage of the secondary cell 10 as detected by the voltmeter 50.

The load 30 includes a wide variety of devices powered by the secondary cell 10; for example, when the system for controlling a secondary cell of the present embodiment is applied to an electric vehicle, the load 30 can be configured from an inverter and a motor. Specifically, when the load 30 is configured from an inverter and a motor, direct-current electric power supplied from the secondary cell 10 is converted into alternating-current electric power by the inverter and supplied to the motor. When the load 30 is configured from an inverter and a motor, a configuration may be adopted to have regenerative electric power generated by the rotation of the motor be converted into direct-current electric power via the inverter and used to charge the secondary cell 10.

Examples of the secondary cell 10 include, e.g., lithium-ion secondary cells and other lithium-based secondary cells. FIG. 2 shows a plan view of the secondary cell 10 of the present embodiment, and FIG. 3 shows a sectional view of the secondary cell 10 across line III-III in FIG. 2.

The secondary cell 10, as shown in FIGS. 2 and 3, is formed from: an electrode layered body 101 having three positive plates 102, seven separators 103, and three negative plates 104; a positive-electrode tab 105 and a negative-electrode tab 106, each of which being connected to the electrode layered body 101; an upper external covering component 107 and a lower external covering component 108 for accommodating and sealing the electrode layered body 101 and the positive- and negative-electrode tabs 105, 106; and a non-aqueous electrolyte (not shown).

The number of positive plates **102**, separators **103**, and negative plates **104** is not particularly limited; the electrode layered body **101** may be configured from one positive plate **102**, three separators **103**, and one negative plate **104**. The number of positive plates **102**, separators **103**, and negative plates **104** may be appropriately selected as necessary.

The positive plates **102** constituting the electrode layered body **101** have positive-electrode-side current collectors **102a** extending to the positive-electrode tab **105**, and positive-electrode active substance layers, each of which being formed on both main surfaces of a portion of the positive-electrode-side current collector **102a**. The positive-electrode-side current collectors **102a** constituting the positive plates **102** can be configured from, e.g., aluminum foil, aluminum alloy foil, copper-titanium foil, stainless foil, or another electrochemically stable metallic foil approximately 20 μm thick.

The positive-electrode active substance layers constituting the positive plates **102** are formed by applying, to the main surfaces of a portion of the positive-electrode-side current collectors **102a**, a blend of positive-electrode active substance, carbon black or another conductive agent, and an aqueous dispersion of polytetrafluoroethylene or polyvinylidene fluoride, or another binder; and subsequently drying and pressing the blended article on the main surfaces.

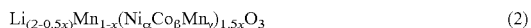
The positive-electrode active substance layers constituting the positive plates **102** in the secondary cell **10** of the present embodiment contain, as a positive-electrode active substance, at least a positive-electrode active substance having the feature that resistance increases in accordance with an increase in the SOC. Such a positive-electrode active substance having the feature that resistance increases in accordance with an increase in the SOC is not particularly limited; examples include a lithium-excess layer electrode material having LiMnO_3 as a base structure and specifically include compounds represented by general formula (1) below. In particular, compounds represented by general formula (1) below have high potential and high capacity, and therefore such compounds are used as positive-electrode active substances. This makes it possible for the secondary cell **10** to have high energy density. Compounds represented by general formula (1) below typically take the form of solid solutions.



(In formula (1) above, $0.1 \leq x \leq 0.5$, and M is $\text{Ni}_\alpha\text{Co}_\beta\text{Mn}_\gamma\text{M}'_\sigma$ (where $0 < \alpha \leq 0.5$, $0 \leq \beta \leq 0.33$, $0 < \gamma \leq 0.5$, $0 \leq \sigma \leq 0.1$, $\alpha + \beta + \gamma + \sigma = 1$, and M' is a metallic element).)

In compounds represented by general formula (1) above, M' may be any metallic element (other than Li, Ni, Co, and Mn) without any particular limitation; M' is preferably at least one element selected from among Fe, V, Ti, Al, and Mg, among which Ti is particularly preferred.

In general formula (1) above, α , β , γ , and σ may be any numbers that satisfy the conditions $0 < \alpha \leq 0.5$, $0 \leq \beta \leq 0.33$, $0 < \gamma \leq 0.5$, $0 \leq \sigma \leq 0.1$, and $\alpha + \beta + \gamma + \sigma = 1$ without any particular limitation; however, preferably $\sigma = 0$. Specifically, compounds represented by general formula (2) below are particularly preferred.



(In formula (2) above, $0.1 \leq x \leq 0.5$, $0 < \alpha \leq 0.5$, $0 \leq \beta \leq 0.33$, $0 < \gamma \leq 0.5$, and $\alpha + \beta + \gamma = 1$.)

The positive-electrode active substance layers may contain positive-electrode active substances other than the positive-electrode active substance described above having the feature that resistance increases in accordance with an increase in the SOC; e.g., lithium nickel oxide (LiNiO_2), lithium manganese

oxide (LiMn_2O_4), lithium cobalt oxide (LiCoO_2), and other lithium compound oxides; and LiFePO_4 , LiMnPO_4 , and the like.

Each of the positive-electrode-side current collectors **102a** constituting the three positive plates **102** are joined to the positive-electrode tab **105**. The positive-electrode tab **105** can be obtained using, e.g., aluminum foil, aluminum alloy foil, steel foil, nickel foil, or the like approximately 0.2 mm thick.

The negative plates **104** constituting the electrode layered body **101** have negative-electrode-side current collectors **104a** extending to the negative-electrode tab **106**, and negative-electrode active substance layers formed on both main surfaces of a part of the negative-electrode-side current collectors **104a**.

The negative-electrode-side current collectors **104a** forming the negative plates **104** are, e.g., nickel foil, copper foil, stainless foil, iron foil, or another electrochemically stable metallic foil approximately 10 μm thick.

The negative-electrode active substance layers forming the negative plates **104** are formed by applying, to the main surfaces of a part of the negative-electrode-side current collectors **104a**, a slurry prepared by adding together a negative-electrode active substance, carbon black or another conductive agent, polyvinylidene fluoride or another binder, and N-methyl-2-pyrrolidone or another solvent; and subsequently drying and pressing the slurry on the main surfaces.

The negative-electrode active substance forming the negative-electrode active substance layers is not particularly limited; a substance containing at least a negative-electrode active substance having, e.g., silicon or carbon as a main element can be used. Examples of negative-electrode active substances having silicon as a main element include silicon as well as silicon oxide and other silicon compounds. Examples of negative-electrode active substances having carbon as a main element include non-graphitizing carbon, graphitizing carbon, graphite, and the like.

In the secondary cell **10** of the present embodiment, the three negative plates **104** are configured so that each of the negative-electrode-side current collectors **104a** forming the negative plates **104** are joined to the single negative-electrode tab **106**. Specifically, in the secondary cell **10** of the present embodiment, each of the negative plates **104** is formed so as to be joined to the single common negative-electrode tab **106**.

The separators **103** in the electrode layered body **101** prevent short-circuiting between the positive plates **102** and the negative plates **104** described above, and may be provided with the capability to hold the electrolyte. The separators **103** are microporous membranes configured from, e.g., polyethylene (PE), polypropylene (PP), or another polyolefin, the separators **103** having a function whereby, when subjected to an overcurrent, the holes in the layers are obstructed and the current is interrupted due to heat generated by the overcurrent.

As shown in FIG. 3, the positive plates **102** and the negative plates **104** are alternately layered with the separators **103** interposed therebetween, and separators **103** are respectively layered on the uppermost and lowermost layers, whereby the electrode layered body **101** is formed.

The electrolyte contained in the secondary cell **10** is a liquid obtained by causing lithium borofluoride (LiBF_4), lithium hexafluorophosphate (LiPF_6), or another lithium salt to dissolve as a solute in an organic liquid solvent. Examples of the organic liquid solvent constituting the electrolyte can include, e.g., propylene carbonate (PC), ethylene carbonate (EC), butylene carbonate (BC), dimethyl carbonate (DMC), ethyl methyl carbonate (EMC), diethyl carbonate (DEC), methyl formate (MF), methyl acetate (MA), methyl propi-

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onate (MP), or another ester-based solvent; alternatively, a blend of the above can be used.

The electrode layered body **101** configured as described above is accommodated and sealed by the upper external covering component **107** and the lower external covering component **108** (sealing means). The upper external covering component **107** and lower external covering component **108** for sealing the electrode layered body **101** are formed from a flexible material such as, e.g., a film made of polyethylene, polypropylene, or another resin; or a plastic/metallic thin-film laminate obtained by laminating both surfaces of a foil made of aluminum or another metal with polyethylene, polypropylene, or another resin. The upper external covering component **107** and the lower external covering component **108** are thermally fused to seal the electrode layered body **101** with the positive-electrode tab **105** and the negative-electrode tab **106** projecting toward the exterior.

A sealing film **109** is disposed on the positive-electrode tab **105** and the negative-electrode tab **106** in order to ensure the upper external covering component **107** and the lower external covering component **108** adhere tightly to the portions at which the upper external covering component **107** and the lower external covering component **108** come into contact with the positive-electrode tab **105** and the negative-electrode tab **106**. The sealing film **109** is not limited by any particular configuration, and can be configured from, e.g., polyethylene, modified polyethylene, polypropylene, modified polypropylene, an ionomer, or another synthetic resin material having exceptional electrolyte resistance and thermal adhesiveness.

The secondary cell **10** of the present embodiment is configured as described above.

Next, the method for controlling charging of the secondary cell **10** of the present embodiment is described. In the present embodiment, control over charging the secondary cell **10** described below is performed by the charge control device **20**. FIG. **4** is a profile showing the charging current and the change in voltage of the secondary cell **10** of the present embodiment when the secondary cell **10** is charged using constant-current/constant-voltage charging.

As shown in FIG. **4**, first, constant-current charging is performed in the present embodiment at an upper-limit voltage V_1 and a set current value A_1 . The upper-limit voltage V_1 is not particularly limited; when, e.g., positive-electrode active substances containing a compound represented by general formula (1) above are used as the positive-electrode materials, the upper-limit voltage V_1 ordinarily is approximately 4.3-4.5V. Similarly, the set current value A_1 is not particularly limited. In the present embodiment, the secondary cell **10** is charged at the set current value A_1 , whereby, as shown in FIG. **4**, the voltage of the secondary cell **10** gradually rises in accordance with an increase in the SOC of the secondary cell **10**; e.g., at time T_1 , the upper-limit voltage V_1 is reached.

Next, in the present embodiment, after the voltage of the secondary cell **10** reaches the upper-limit voltage V_1 , constant-voltage charging is performed at the upper-limit voltage V_1 . In the present embodiment, when constant-voltage charging is performed at the upper-limit voltage V_1 , the charging current decays as the SOC of the secondary cell **10** increases, with the voltage of the secondary cell **10** being kept at the upper-limit voltage V_1 , as shown in FIG. **4**. In the present embodiment, when the charging current decays and decreases to the cutoff current value A_2 , charging of the secondary cell **10** is terminated. The above describes how charging the secondary cell **10** is controlled in the present embodiment.

Next, the method for setting the cutoff current value A_2 is described. First, when the cutoff current value A_2 is to be set

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in the present embodiment, a threshold coefficient X is calculated according to formula (3) below based on a cell resistance R_1 of the secondary cell **10** at a target SOC (units: Ω), the set current value A_1 during constant-current charging described above (units: A), and the upper-limit voltage V_1 (units: V).

$$\text{Threshold coefficient } X = (\text{cell resistance } R_1 \text{ of secondary cell in target SOC } [\Omega] \times \text{set current value } A_1 [A]) / \text{upper-limit voltage } V_1 [V] \quad (3)$$

What is meant by “target SOC” in “cell resistance R_1 of the secondary cell at a target SOC” is the SOC when, e.g., the secondary cell **10** is charged to the upper-limit voltage V_1 at a rate of 0.1 C. Specifically, if, e.g., when the secondary cell **10** is charged at 0.1 C, the SOC at the point in time when the voltage of the secondary cell **10** reaches 4.3V is 50%, then the target SOC upon setting the upper-limit voltage V_1 at 4.3V is 50%. Similarly, if the SOC at the point in time when the voltage of the secondary cell **10** reaches 4.4V is 70%, then the target SOC upon setting the upper-limit voltage V_1 at 4.4V is 70%, and if the SOC at the point in time when the voltage of the secondary cell **10** reaches 4.5V is 90%, then the target SOC upon setting the upper-limit voltage V_1 at 4.5V is 90%. The “cell resistance R_1 of the secondary cell at a target SOC,” then, can be obtained by measuring the cell resistance when the secondary cell **10** is to be charged to the target SOC.

The relationship between the target SOC and the upper-limit voltage V_1 typically changes depending on the type of positive- and negative-electrode active substances forming the secondary cell **10**, the type of non-aqueous electrolyte, the balance between the positive and negative electrodes, and other factors. However, secondary cells **10** manufactured with the same design (specifically, secondary cells **10** using the same positive-electrode active substances, negative-electrode active substances, and non-aqueous electrolytes and having the same balance between positive and negative electrodes) will exhibit the same relationship between the target SOC and the upper-limit voltage V_1 . Therefore, in the present embodiment, it is preferable, e.g., to use different secondary cells **10** manufactured with the same design; to measure, in advance, the relationship between the cell resistance R_1 of the secondary cell at a target SOC and the upper-limit voltage V_1 ; and to store the resulting measurements in the charge control device **20**.

The method for measuring the “cell resistance R_1 of the secondary cell at a target SOC” is not particularly limited; measurement may be performed by, e.g., measuring the alternating-current impedance in a secondary cell **10** charged to the target SOC.

The threshold coefficient X is calculated according to formula (3) above based on the cell resistance R_1 of the secondary cell **10** at a target SOC, the set current value A_1 during constant-current charging, and the upper-limit voltage V_1 . Specifically, when the cell resistance R_1 of the secondary cell **10** at a target SOC is 7.5Ω , the set current value A_1 is 0.035 A , and the upper-limit voltage V_1 is 4.5 V , the threshold coefficient X is 0.0583 ($X = (7.5 \times 0.035) / 4.5$).

Next, in the present embodiment, the threshold coefficient X thus calculated is used to set the cutoff current value A_2 . Specifically, the cutoff current value A_2 is set so as to comply with formula (4) below, based on the threshold coefficient X and the set current value A_1 during constant-current charging.

$$\text{Cutoff current value } A_2 \geq \text{set current value } A_1 \times \text{threshold coefficient } X \quad (4)$$

Specifically, the cutoff current value A_2 is set to a value equal to or greater than the product ($A_1 \times X$) of the set current value A_1 and the threshold coefficient X .

Rearranging formula (4) gives formula (5) below.

$$\text{Cutoff current value } A_2 / \text{set current value } A_1 \geq \text{threshold coefficient } X \quad (5)$$

Therefore, in the present embodiment, the cutoff current value A_2 can be set so that the ratio (A_2/A_1) of the cutoff current value A_2 with respect to the set current value A_1 is equal to or greater than the threshold coefficient X .

In the present embodiment, the cutoff current value A_2 can be any value equal to or greater than the product ($A_1 \times X$) of the set current value A_1 and the threshold coefficient X (and less than the set current value A_1). If, e.g., the cutoff current value A_2 is set to a low value, the secondary cell 10 will have a deeper charge; therefore, the post-charge SOC of the secondary cell 10 can be set relatively high. On the other hand, if the cutoff current value A_2 is set to a high value, the post-charge SOC of the secondary cell 10 becomes relatively low, but the time needed for charging decreases.

In the present embodiment, the cutoff current value A_2 calculated in this manner is used to perform constant-current/constant-voltage charging on the secondary cell 10; at the point in the constant-voltage charging when the charging current decreases to the cutoff current value A_2 , the charging for the secondary cell 10 is terminated.

In the present embodiment, when the secondary cell 10 in which there is used, as a positive-electrode material, a positive-electrode active substance having the feature that resistance increases in accordance with an increase in the SOC is to be charged using constant-current/constant-voltage charging, the cutoff current value A_2 for terminating the constant-voltage charging is set equal to or greater than the product ($A_1 \times X$) of the set current value A_1 during constant-current charging and the threshold coefficient X calculated from the cell resistance R_1 of the secondary cell at a target SOC, the set current value A_1 during constant-current charging, and the upper-limit voltage of charging. Therefore, according to the present embodiment, side reactions associated with excessive current impressing can be suppressed when such a secondary cell 10 is to be charged. This makes it possible to improve the charge/discharge efficiency of the secondary cell 10, resulting in improved cycling characteristics.

In particular, when a positive-electrode active substance having the feature that resistance increases in accordance with an increase in the SOC is used as a positive-electrode material, the positive/negative electrode potential changes in the end stage of charging, and non-aqueous-electrolyte decomposition, lithium deposition, and other side reactions occur, resulting in diminished cycling characteristics. However, according to the present embodiment, the cutoff current value A_2 is set equal to or greater than the product ($A_1 \times X$) of the set current value A_1 during constant-current charging and the threshold coefficient X calculated from the cell resistance R_1 of the secondary cell at a target SOC, the set current value A_1 during constant-current charging, and the upper-limit voltage of charging; and, when the charging current in the constant-voltage charging decays to the cutoff current value A_2 , charging of the secondary cell 10 is terminated, allowing such problems to be effectively resolved.

In the embodiment described above, the control device 20 corresponds to each of constant-current charging means or device, constant-voltage charging means or device, and charge-stopping means or device in the present invention, as it pertains to the charge control device for a secondary cell.

An embodiment of the present invention is described above; however, this embodiment was described in order to facilitate an understanding of the present invention, and is not particularly limited to the present invention. Therefore, each

of the elements disclosed in the embodiment above encompasses all design variants and equivalents belonging to the technical scope of the present invention.

WORKING EXAMPLES

The present invention is described in more detail below based on working examples; however, these examples are not particularly limited to the present invention.

Production of Positive Electrode

Graphite powder, acetylene black (conductive auxiliary agent), and PVDF (binder) were combined so as to have a mass ratio of 90:5:5. N-methyl-2-pyrrolidone was introduced to the resulting combination as a solvent, and these components were mixed to produce negative-electrode slurry. The negative-electrode slurry was then applied to a copper foil (current collector) so as to have a dried thickness of 70 μm . The solvent was dried, whereupon drying was performed in a vacuum for 24 hours, resulting in a negative electrode.

$\text{Li}_{1.85}\text{Ni}_{0.18}\text{Co}_{0.10}\text{Mn}_{0.87}\text{O}_3$ (positive-electrode active substance; in formula (1) above, $x=0.3$, $\alpha=0.40$, $\beta=0.22$, $\gamma=0.38$, and $\sigma=0$), acetylene black (conductive auxiliary agent), and PVDF (binder) were combined so as to have a mass ratio of 90:5:5. N-methyl-2-pyrrolidone was introduced to the resulting combination as a solvent, and these components were mixed to produce positive-electrode slurry. The positive-electrode slurry was then applied to an aluminum foil (current collector) so as to have a dried thickness of 50 μm . The solvent was dried, whereupon drying was performed in a vacuum for 24 hours, resulting in a positive electrode.

Production of Lithium-Ion Secondary Cell

Next, the negative electrode and the positive electrode obtained as described above were set in opposition to each other, and a 20- μm -thick separator made of polyolefin was disposed therebetween, resulting in a layered body comprising a negative electrode, a separator, and a positive electrode. The resulting layered body comprising a negative electrode, a separator, and a positive electrode was placed inside a laminate cell made of aluminum, an electrolyte was injected into the cell, and then the layered body was hermetically sealed, resulting in a lithium-ion secondary cell. 1M LiPF_6 in ethylene carbonate (EC):diethyl carbonate (DEC) (1:2 (by volume)) was used as the electrolyte.

Activation of Cell and Measurement of Cell Capacitance

The lithium-ion secondary cell obtained as described above was charged to 4.5V, using a constant-current charge method, at 0.1 C under a 30° C. atmosphere; and then, after being left inactive for ten minutes, was similarly discharged at a constant current of 0.1 C to 2.0V. The constant-current charging at 0.1 C, ten-minute period of inactivity, and constant-current discharging at 0.1 C (cutoff voltage: 2.0V) were then repeatedly performed with the charging upper-limit voltage varied to each of 4.6V, 4.7V, and 4.8V, whereby the cell was activated. In the present example, the discharge capacity when the charging upper-limit voltage was set to 4.8V was 35 mAh, and this value was used as the cell capacitance (specifically, the capacitance when SOC=100%).

Measurement of Cell Resistance

The lithium-ion secondary cell, having been activated as described above, was charged so that SOC=10%, whereupon impressing of current was stopped and the lithium-ion secondary cell was discharged for two hours. Alternating-current impedance measurements were then taken at a voltage width of 10 mV and a frequency of 10 mHz-1 MHz in the lithium-ion secondary cell that was charged so that SOC=10%, whereby the cell resistance was measured.

Then, the same operations were performed at increments of SOC 10% up to SOC=100%, whereby the cell resistance was measured in each of the following states: SOC=10%, 20%, 30%, 40%, 50%, 60%, 70%, 80%, 90%, and 100%.

Charge/Discharge Cycle Testing

Next, a plurality of lithium-ion secondary cells manufactured and activated in the same manner as described above were prepared, and cycle testing was performed under the charge/discharge conditions below.

Working Example 1

A lithium-ion secondary cell was charged, using a constant-current/constant-voltage charging method, for two hours under a 30° C. atmosphere with a set current value A_1 of 35 mA (1 C), a cutoff current value A_2 of the constant-voltage charging of 17.5 mA ($A_2/A_1=0.5$), and an upper-limit voltage V_1 of 4.5V; left inactive for 10 minutes; and then discharged at a constant current of 35 mA (1 C) to 2V. A cycle test in which this process constituted one cycle was performed for 100 cycles. In working example 1, the target SOC was 90%, and the cell resistance at this time measured according to the method described above was 7.5Ω. Therefore, the threshold coefficient X was set to 0.0583 ($X=(7.5 \times 0.035)/4.5$). Because $A_2/A_1=0.5$, working example 1 satisfies the condition $A_2/A_1 \geq X$. Table 1 shows the capacity retention at the 100th cycle (=discharge capacity at 100th cycle/discharge capacity at 1st cycle $\times 100(\%)$).

Working Example 2

100 cycles of the cycle test were performed in the same manner as in working example 1, other than that the cutoff current value A_2 of the constant-voltage charging was changed to 8.75 mA ($A_2/A_1=0.25$). In working example 2, the threshold coefficient X was set to 0.0583, the same as in working example 1. Because $A_2/A_1=0.25$, working example 2 satisfies the condition $A_2/A_1 \geq X$. Table 1 shows the capacity retention at the 100th cycle.

Working Example 3

100 cycles of the cycle test were performed in the same manner as in working example 1, other than that the cutoff current value A_2 of the constant-voltage charging was changed to 3.5 mA ($A_2/A_1=0.1$). In working example 3, the threshold coefficient X was set to 0.0583, the same as in working example 1. Because $A_2/A_1=0.1$, working example 3 satisfies the condition $A_2/A_1 \geq X$. Table 1 shows the capacity retention at the 100th cycle.

Comparative Example 1

100 cycles of the cycle test were performed in the same manner as in working example 1, other than that the cutoff current value A_2 of the constant-voltage charging was changed to 1.75 mA ($A_2/A_1=0.05$). In comparative example 1, the threshold coefficient X was set to 0.0583, the same as in working example 1. However, because $A_2/A_1=0.05$, comparative example 1 does not satisfy the condition $A_2/A_1 \geq X$. Table 1 shows the capacity retention at the 100th cycle.

Comparative Example 2

100 cycles of the cycle test were performed in the same manner as in working example 1, other than that the cutoff current value A_2 of the constant-voltage charging was

changed to 1.05 mA ($A_2/A_1=0.03$). In comparative example 2, the threshold coefficient X was set to 0.0583, the same as in working example 1. However, because $A_2/A_1=0.03$, comparative example 2 does not satisfy the condition $A_2/A_1 \geq X$. Table 1 shows the capacity retention at the 100th cycle.

Comparative Example 3

100 cycles of the cycle test were performed in the same manner as in working example 1, other than that the cutoff current value A_2 of the constant-voltage charging was changed to 0.525 mA ($A_2/A_1=0.015$). In comparative example 3, the threshold coefficient X was set to 0.0583, the same as in working example 1. However, because $A_2/A_1=0.015$, comparative example 3 does not satisfy the condition $A_2/A_1 \geq X$. Table 1 shows the capacity retention at the 100th cycle.

Comparative Example 4

A lithium-ion secondary cell was charged, using a constant-current/constant-voltage charging method, for two hours under a 30° C. atmosphere with a set current value A_1 of 35 mA (1 C) and an upper-limit voltage V_1 of 4.5V; left inactive for 10 minutes; and then discharged at a constant current of 35 mA (1 C) to 2V. A cycle test in which this process constituted one cycle was performed for 100 cycles. Specifically, in comparative example 4, constant-current/constant-voltage charging was performed without a cutoff current value A_2 being set. Table 1 shows the capacity retention at the 100th cycle.

TABLE 1

	Value of A_2/A_1	Capacitance retention rate at 100th cycle
Working example 1	0.5	92%
Working example 2	0.25	90%
Working example 3	0.1	90%
Comparative example 1	0.05	83%
Comparative example 2	0.03	76%
Comparative example 3	0.015	67%
Comparative example 4	0 (cutoff current not set)	41%

Evaluation of Working Examples 1-3 and Comparative Examples 1-4

Table 1 combines the results for working examples 1-3 and comparative examples 1-4 in which the cycle tests were performed with an upper-limit voltage V_1 of 4.5.

As shown in Table 1, the results obtained for working examples 1-3, in which a current value satisfying the condition $A_2/A_1 \geq X$ was adopted as the cutoff current value A_2 , were all excellent in that the capacity retention at the 100th cycle was 90% or higher. However, comparative examples 1-3, in which a current value not satisfying the condition $A_2/A_1 \geq X$ was set as the cutoff current value A_2 , and comparative example 4, in which no cutoff current value A_2 was set, all had inferior cycling characteristics in that the capacity retention at the 100th cycle was less than 80%. FIG. 5 shows a graph depicting the relationship between the value of A_2/A_1 and the capacity retention at the 100th cycle for working examples 1-3 and comparative examples 1-4. As can be ascertained from FIG. 5, when the condition $A_2/A_1 \geq X$ is satisfied (specifically, when $A_2/A_1 \geq 0.0583$), the capacity retention at

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the 100th cycle is a high value of 90% or higher, and that stable results can be obtained. In FIG. 5, comparative example 4, in which no cutoff current value A_2 was set, is displayed at $A_2/A_1=0$.

Working Example 4

A lithium-ion secondary cell was charged, using a constant-current/constant-voltage charging method, for two hours under a 30° C. atmosphere with a set current value A_1 of 35 mA (1 C), a cutoff current value A_2 of the constant-voltage charging of 1.75 mA ($A_2/A_1=0.05$), and an upper-limit voltage V_1 of 4.4V; left inactive for 10 minutes; and then discharged at a constant current of 35 mA (1 C) to 2V. A cycle test in which this process constituted one cycle was performed for 100 cycles. In working example 4, the target SOC was 70%, and the cell resistance at this time measured according to the method described above was 4Ω. Therefore, the threshold coefficient X was set to 0.0318 ($X=(4 \times 0.035)/4.4$). Because $A_2/A_1=0.05$, working example 4 satisfies the condition $A_2/A_1 \geq X$. Table 2 shows the capacity retention at the 100th cycle.

Comparative Example 5

A lithium-ion secondary cell was charged, using a constant-current/constant-voltage charging method, for two hours under a 30° C. atmosphere with a set current value A_1 of 35 mA (1 C) and an upper-limit voltage V_1 of 4.4V; left inactive for 10 minutes; and then discharged at a constant current of 35 mA (1 C) to 2V. A cycle test in which this process constituted one cycle was performed for 100 cycles. Specifically, in comparative example 5, constant-current/constant-voltage charging was performed without a cutoff current value A_2 being set. Table 2 shows the capacity retention at the 100th cycle.

TABLE 2

	Value of A_2/A_1	Capacitance retention rate at 100th cycle
Working example 4	0.05	93%
Comparative example 5	0	38%
(cutoff current not set)		

Evaluation of Working Example 4 and Comparative Example 5

Table 2 combines the results for working example 4 and comparative example 5 in which the cycle tests were performed with an upper-limit voltage V_1 of 4.4.

As shown in Table 2, the results obtained for working example 4, in which a current value satisfying the condition $A_2/A_1 \geq X$ was adopted as the cutoff current value A_2 , were excellent in that the capacity retention at the 100th cycle was 90% or higher. However, comparative example 5, in which no cutoff current value A_2 was set, had extremely inferior cycling characteristics in that the capacity retention at the 100th cycle was less than 40%.

Working Example 5

A lithium-ion secondary cell was charged, using a constant-current/constant-voltage charging method, for two hours under a 30° C. atmosphere with a set current value A_1 of 35 mA (1 C), a cutoff current value A_2 of the constant-voltage

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charging of 1.75 mA ($A_2/A_1=0.05$), and an upper-limit voltage V_1 of 4.3V; left inactive for 10 minutes; and then discharged at a constant current of 35 mA (1 C) to 2V. A cycle test in which this process constituted one cycle was performed for 100 cycles. In working example 5, the target SOC was 50%, and the cell resistance at this time measured according to the method described above was 4Ω. Therefore, the threshold coefficient X was set to 0.0325 ($X=(4 \times 0.035)/4.3$). Because $A_2/A_1=0.05$, working example 5 satisfies the condition $A_2/A_1 \geq X$. Table 3 shows the capacity retention at the 100th cycle.

Comparative Example 6

A lithium-ion secondary cell was charged, using a constant-current/constant-voltage charging method, for two hours under a 30° C. atmosphere with a set current value A_1 of 35 mA (1 C) and an upper-limit voltage V_1 of 4.4V; left inactive for 10 minutes; and then discharged at a constant current of 35 mA (1 C) to 2V. A cycle test in which this process constituted one cycle was performed for 100 cycles. Specifically, in comparative example 6, constant-current/constant-voltage charging was performed without a cutoff current value A_2 being set. Table 3 shows the capacity retention at the 100th cycle.

TABLE 3

	Value of A_2/A_1	Capacitance retention rate at 100th cycle
Working example 5	0.05	93%
Comparative example 6	0	65%
(cutoff current not set)		

Evaluation of Working Example 5 and Comparative Example 6

Table 3 combines the results for working example 5 and comparative example 6 in which the cycle tests were performed with an upper-limit voltage V_1 of 4.3.

As shown in Table 3, the results obtained for working example 5, in which a current value satisfying the condition $A_2/A_1 \geq X$ was adopted as the cutoff current value A_2 , were excellent in that the capacity retention at the 100th cycle was 90% or higher. However, comparative example 6, in which no cutoff current value A_2 was set, had inferior cycling characteristics in that the capacity retention at the 100th cycle was less than 70%.

The invention claimed is:

1. A method for controlling charging of a secondary cell including a positive electrode containing a positive-electrode active substance having a resistance that increases in accordance with an increase in SOC, a negative electrode, and a non-aqueous electrolyte,

the method comprising:

performing constant-current charging at a set current value A_1 to a prescribed upper-limit voltage V_1 ;

performing constant-voltage charging at the upper-limit voltage V_1 after the upper-limit voltage V_1 has been reached; and

terminating charging of the secondary cell when a charging current in the constant-voltage charging has decreased to a cutoff current value, A_2

the cutoff current value A_2 being set to a current value that complies with the relationships in formulas (I) and (II),

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- Cutoff current value $A_2 \geq$ set current value $A_1 \times X$ (I)
- $X = (\text{cell resistance } R_1 \text{ of secondary cell at target SOC } [\Omega] \times \text{set current value } A_1 \text{ [A]}) / \text{upper-limit voltage } V_1 \text{ [V]}$ (II). 5
2. The method of claim 1, wherein the positive-electrode active substance contains a compound represented by formula (III)
- $\text{Li}_{(2-0.5x)}\text{Mn}_{1-x}\text{M}_{1.5x}\text{O}_3$ (III) 10
- (In formula (III), $0.1 \leq x \leq 0.5$, and M is $\text{Ni}_\alpha\text{Co}_\beta\text{Mn}_\gamma\text{M}'_\sigma$ (where $0 < \alpha \leq 0.5$, $0 \leq \beta \leq 0.33$, $0 < \gamma \leq 0.5$, $0 \leq \sigma \leq 0.1$, $\alpha + \beta + \gamma + \sigma = 1$, and M' is a metallic element)). 15
3. The method of claim 1, wherein the negative electrode contains a negative-electrode active substance having at least one of silicon and carbon as a main element.
4. A charge control device for controlling charging of a secondary cell including a positive electrode containing a positive-electrode active substance having a resistance that increases in accordance with an increase in SOC, a negative electrode, and a non-aqueous electrolyte, the charge control device comprising: 20

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- a constant-current charging device configured to perform constant-current charging at a set current value A_1 to a prescribed upper-limit voltage V_1 ;
- a constant-voltage charging device configured to perform constant-voltage charging at the upper-limit voltage V_1 after the upper-limit voltage V_1 has been reached; and
- a charging-stopping device configured to determine whether a charging current in the constant-voltage charging has decreased to a cutoff current value A_2 , and, when the charging-stopping device has determined that the charging current has decreased to the cutoff current value A_2 , stopping charging of the secondary cell
- the charge-stopping device configured to set the cutoff current value A_2 at a current value that complies with the relationships in formulas (I) and (II)
- Cutoff current value $A_2 \geq$ set current value $A_1 \times X$ (I)
- $X = (\text{cell resistance } R_1 \text{ } [\Omega] \text{ of secondary cell at target SOC} \times \text{set current value } A_1 \text{ [A]}) / \text{upper-limit voltage } V_1 \text{ [V]}$ (II).
5. The method of claim 2, wherein the negative electrode contains a negative-electrode active substance having at least one of silicon and carbon as a main element.

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